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(71) Applicant: GEOCHEMICAL CORPORATION [US/ US]; 162 Spencer Place, Ridgewood, NJ 07450 (US).

(71)(72) Applicant and Inventor: CLARKE, William, J. [US/ US]; 162 Spencer Place, Ridgewood, NJ 07450 (US).

(74) Agents: HANDELMAN, Joseph. H. et al.; Ladas & Parry, 26 West 61 Street, New York, NY 10023 (US).

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(54) Title: GROUTING COMPOSITION COMPRISING SLAG

(57) Abstract

A grouting composition, a method of grouting and a formation so grouted. The composition comprises a very small particle size slag, an equal or greater weight of water and the optional components cement, alkali silicate, anionic dispersant, a source of orthophosphate ions, sodium hydroxide and sodium carbonate. The grout is particularly adapted to the treatment of "tight" or low permeability formations being low in viscosity and having controllable set time and hardening time as well as high strength upon hardening. Being based on slag, a byproduct, the grout is economical.

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GROUTING COMPOSITION COMPRISING SLAG

This invention provides an improved and economical grouting composition or grout, which is suitable for stabilizing and strengthening earthen foundations, especially those employed in permanent structures such as buildings, tunnels and dams. The composition and the methods and procedures for its use are particularly suited to curtain grouting dams, sealing off underground streams, isolating buried containers, reinforcing and protecting building foundations, protecting excavations, especially those used in the construction of subways, breakwater and bridge piers, and other applications for sealing or closing of passageways to the transport of fluids therethrough. The grouting composition of this invention is particularly fluid having an equal or greater weight of water than cementitious material.

Although relatively low in cost, the composition is surprisingly efficient, especially in a one component injection process. The composition has low viscosity, predictable set time, high ultimate strength and producing a sealed formation of very low permeability to fluids, particularly aqueous systems. This composition, comprising a suspension of cementitious solids, penetrates formations with a facility hithertofor considered characteristic of solution grout compositions but it develops the high formation strength characteristic of the best

suspension compositions.

grouted.

Being free of organic chemicals except for a small amount of dispersant, by its nature substantive to the matter in the formation being grouted, the grout of this invention is non-polluting. This grout does not contribute to underground water or soil pollution in formations in which the grout is used. The non-polluting aspect of the grout is particularly important in environmental improvement applications.

In these applications the very low permeability, to water and other fluids, of the grouted substrate is a most significant property resulting from the use of this invention. Examples of environmental improvement applications are the sealing of reservoirs for storage of waste materials, including toxic and radioactive

A major application of the grout of this invention is rehabilitation of structures such as buildings, bridges, dams and subways. A particularly important application is the regrouting of dam curtains, especially grouted curtains which have shown evidence of deterioration. Such deterioration is often evidenced by increased seepage and even eventual

wastes, and the sealing of formations where unwanted

seepage into underground aquifers is occuring.

The "one component" or single fluid grout of this invention does not have the difficultly controlled on site mixing and proportioning operations inherent in other types such as "two component"

30 systems. The single fluid grout of this invention contains the necessary ingredients to produce the final high strength solidified state. The single fluid grout is programmed, by composition, to have a set time high enough to facilitate ease of injection into a formation to be grouted yet low enough to prevent its dilution or loss from the formation

However, in certain applications of this invention very fast set times, such as five to thirty minutes, are desired. These are readily achieved by two component grout compositions which employ the ultrafine cementitious materials in one component and silicates in a second component.

In its various aspects this invention concerns: (1) a composition adapted for use in grouting, (2) a method of grouting comprising preparing the composition and injecting the composition into a formation and (3) a formation grouted by the composition.

The grouting composition of this invention comprises an ultrafine slag and water. The ultrafine 15 slag is a cementitious material (CM) having a specific surface area greater than 9500, preferably greater than 11000 and most preferably greater than 13000 square centimeters per gram and, by weight, less than 3%, preferably less than 1.5% and most preferably no 20 particles larger than 7.8 micrometers (um) (the surface area and particle size are measured by the use of the MicrotracR instrument further described hereinbelow). An optional cementitious material in the composition is cement having a specific surface area 25 greater than 7000, preferably greater than 8000 and most preferably greater than 9000 square centimeters per gram with, by weight, less than 16%, preferably less than 7% and most preferably less than 3% of particles larger than 7.8 micrometers as measured by 30 the $Microtrac^R$ instrument. The preferred cement is portland cement.

The cementitious material is at least 40%, preferably 60%, more preferably 80% and most preferably 100% slag by weight.

The water to cementitious material weight ratio is 1:1 to 12:1 or greater, preferably 1.5:1 to 7:1 and most preferably 2:1 to 3:1 i.e. there is

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always at least as much water as cementitious material and sufficient water to form a flowable grout.

For adapting to specific use requirements or operating conditions, additional materials in the grout are as follows, amounts being in weight percent solids based on the weight of cementitious material unless otherwise noted:

- (3) An effective amount of dispersant to disperse the cementitious material, preferably up to 10 2%, and most preferably 0.2% to 0.5%.
 - (4) NaOH, Na₂CO₃ or a mixture of these up to about 15%, preferably up to 10% and most preferably about 3% to 8% of the weight of slag plus about 0% to 4% of the weight of cement.
- 15 (5) An alkali silicate up to about 150%, preferably 30% to 75%.
 - (6) When an alkali silicate is present, up to about 10%, preferably about 1% to 5% of orthophosphate ion is also provided.
 - (7) Other optional ingredients comprise materials such as inert filler material as discussed hereinbelow.

In certain applications combinations of slag and cement have the most appropriate properties, in others, alkali silicate is desired as a third large component of the composition, in still others, slag and silicate give the desired properties. One of the advantages of this invention is the ease with which the composition can be altered to adapt to set time, strength, ultimate permeability of the grouted formation and other requirements for the specific job. Combinations of portland cement and slag in the composition furnish grouts having an outstanding combination of permeation, set time and strength; even modest amounts of cement in such compositions produce grout having a controllable desirable set time. Combinations of these ingredients lead to optimum

strength, particularly unconfined compressive strength.

The slag employed in this invention is a byproduct obtained in the blast furnace process of 5 smelting iron ore to produce iron. The slag is produced by a conventional process, outlined below, utilizing grinding and separating steps well known in the art. In the blast furnace, slag is generated, typically at about 1550 to 1650°C., from other 10 minerals in the iron ore, coke ash and the limestone used as a flux. This slag is skimmed from above the iron, granulated by treatment with water and ground. Typically the granulation employs high pressure water sprays followed by immersion of the slag in a pit of 15 water to produce a slag having a high glass content. However produced, it is desired that the slag used in this invention have a high glass content, preferably above about 95% glass. It is also preferred that the water content be low, preferably below about 15%. Typical dry composition range and two examples of slag 20 are (in weight percent): Composition SiO₂ Al₂O₃ CaO MgO FeO S Mn₂O₃8-18 35-50 0-15 0-1 0-2 0-2 Usual Range 30-40 0.3 0.5 < 0.1 3.5 43.3 16.0 Typical 36.4

35.5 Typical The slag used in this invention has a specific surface area over 9500 cm²/gm, desirably over 11000 cm^2/gm and most preferably over 13000 cm^2/gm .

11.4

39.4

0.2

11.9

0.2

The high glass content, finely divided slag yields a grout having excellent sulfate resistance and 30 readily controlled setting and hardening times in a relatively inexpensive formulation. High sulfate resistance is particularly important with regard to long term stability of the grouted formation. Grout failures after five or ten years have been attributed 35 to sulfate attack. The slag-rich compositions of the present invention have long term stability in

formation and are also useful in regrouting formations in which a previous grout has exhibited signs of failure. The grout is particularly inexpensive because of the comparatively low cost of slag, a byproduct of iron production.

High level of penetration into a tight formation associated with high permanance renders the instant composition valuable in many applications. One example is in radioactive waste storage in shafts deep in rock formations where the instant grout composition is employed to permeate the rock around the waste containers and thus furnish a secondary barrier to both intrusion of ground water and leaking of waste from the containers.

- Portland cement is a hydraulic cement consisting mainly of calcined silicates of calcium. Five types of portland cement (I through V) are generally recognized, their manufacture, properties and characteristics are described in detail by J.F.
- Young, ACI (American Concrete Institute, Detroit, Michigan) Education Bulletin No. E3-83, pages 1 through 8, herein incorporated by reference.

It is a novel feature of this invention that cementitious material having the fine particle size and distribution described is formulated in a composition producing a high strength grout having a readily controlled set time.

While portland cement and/or slag are
preferably used without inert filler material, in some
instances it may be desirable to incorporate limited
amounts of other solids meeting the particle size
specifications specified for slag hereinabove. Such
other solids are clay, bentonite, kaolin, vermiculite,
limestone, silica flour, silica fume and other well
known inert solids. The amount of these employed is
to be minor so as not to reduce the strength of the
set grout below desired values such as those in the

examples of this invention hereinbelow.

The alkali silicate is preferably an aqueous colloidal suspension of an alkali salt of silicic Throughout the specifications and claims, the term "alkali silicate" is intended to refer to alkali metal silicates having an alkali metal oxide:silicon dioxide weight ratio within the range of from about 1.0:3.0 to 1.0:4.0, notably sodium silicates having a sodium oxide:silicon dioxide weight ratio within the range of from about 1:3.0 to 1:4.0, preferably, about 10 1:3.0 to 1:3.5. An especially preferred material is an aqueous sodium silicate having a sodium oxide:silicon dioxide ratio of about 1:3.2 to 1:3.3 and a Na₂O:SiO₂ solids content of about 35 to 45%. The term "alkali metal", as used in the specification. 15 and claims is intended to refer to the various alkali metals, i.e., sodium, potassium, lithium, rubidium, cesium and mixtures thereof. Silicates of potassium and sodium are more generally available. their lower cost and availability, sodium silicates 20 are more widely used and therefore are preferred in the practice of the present invention. Particular reference hereinafter is made to such silicates. alkali silicate is especially useful in the preparation of low cost compositions having a very 25 fast set time for uses such as underground water control.

The dispersant is selected from the wellknown group of materials used to disperse solids,
preferably water-wetable solids, in aqueous systems.
The dispersant serves to aid the penetration of water
into the finely divided solids to produce a suspension
of unassociated or unagglomerated particles. These
dispersants are generally negatively charged or
anionic electrolytes especially polyelectrolytes, such
as polycarboxylates and polysulphonates. Examples
include sodium or ammonium salts of polymethacrylic

acid, diisobutylene-maleic anhydride copolymer, copolymers of acrylic, methacrylic and maleic acids, organic phosphates, sulfonated naphthalene formaldehyde condensates, sulfonated natural products and other sulfonated polymer condensates.

The orthophosphate ion is any of the orthophosphate ions generally existing in equilibrium with each other in aqueous solution, PO4 TT, HPO4 TT and H₂PO₄. In highly alkaline systems it is expected 10 that POA predominates. Two of the most available sources of orthophosphate ion are orthophosphoric acid, H3PO4 and sodium phosphate decahydrate, Na₃PO₄.10H₂O. Phosphoric acid is commonly the commercial grade 85%, by volume, with water. Other phosphate such as the polyorthophosphates, may be 15 employed as the source of the orthophosphate ion. orthophosphate ion is useful in control of the set time, especially in grout compositions comprising alkali silicates, by delaying setting.

The NaOH is commercially available in solid and aqueous solution grades, the Na₂CO₃ is usually a solid grade, each serves to control the rate of hardening of the composition, particularly in formulations high in slag. Thus, NaOH and/or Na₂CO₃ may be regarded as a set time catalyst which has little if any effect on other properties of the grout, however overdosage will decrease strength. Favored are formulations in which the Na₂CO₃ is less than equal to NaOH by weight.

The importance of controlled rapid set time is seen, for example, in dam curtain rehabilitation. In a typical case water is moving through the curtain, at perhaps one or two feet per hour, and the grout injection holes in the dam curtain are on ten foot centers. Thus the set time of the grout must be less than five hours, preferably in the half hour to five hour range and must be predictable. This invention

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teaches predictability in both set time and permeation, particularly in tight formations previously grouted where degraded cement results in increased seepage. The ultrafine slag permeates the degraded cement, hardens and reduces seepage of the rehabilitated dam grout curtain.

In general, portland cement contributes to decreasing the set time and increasing the strength of a formulation. Slag, without cement has longer set 10 times but this is controlled by the use of NaOH and Na₂CO₃. Higher than optimum levels of NaOH and Na₂CO₃ tend to decrease the final strength and increase the cost of the formulation. When alkali silicate is employed, optimization of the slag-cement-silicate 15 ratios can result in a formulation with too fast a set. This low set time can than be increased, with little effect on strength, by the addition of phosphate ions. In all formulations the viscosity of the grout is readily controlled by the amount of water present. Thus in each case the invention teaches the balancing of the composition variables needed to give the optimum grout for the given application.

Another novel feature of this invention is based on the finding that NaOH and Na₂CO₃ are unexpectedly efficient and effective in controlling 25 the set time of mixtures of slag and alkali silicate. When cement is used to control the set time of admixtures with slag and silicate there is the difficulty of mixing the appropriate ratio of the 30 three large components in the field and a second difficulty is that flocculation of the system occur before setting. The latter leads to poor permeation of the formation being grouted. The use of NaOH, Na₂CO₃ or a mixture of these speed the set time to a suitable range, employing a small amount, which is easily measured and added in the field, and additionally produces no separation or flocculation in the grout before an end of the set time can be obtained by the addition of phosphate ion as in other alkali silicate systems, if desired.

pedagogical aid in anning the invention, it is clear that since the slag is essentially free of available calcium ion the formation of calcium silicate does not occur. The formation of the calcium salt is likely to be responsible for the separation or flocculation observed when cement is present. The lack of exchangeable calcium in slag is also thought to contribute to the effectiveness of the NaOH and Na₂CO₃ in acceleration of the setting of silicate-slag systems.

15 The following examples will further illustrate the invention but are not intended to limit it in any way. All parts and percentages are by weight and all temperatures in degrees Centigrade unless otherwise noted.

20 Particle Size of Cementitious Materials The particle size of the materials is determined by a laser light scattering particle size distribution apparatus identified as a MicrotracR particle size analyzer (Leeds and Northrup Instruments, North Wales, Pennsylvania 19454). 25 Measurements are made in the 0.7 to 125 millimicron (um) range and are presented in terms of the volume of particles in size channels having a size ratio of approximately 1.4 between channels. Table I presents results of the average of three runs given as the 30 percent larger than the stated size for each channel limit and the calculated volume surface area of the sample in square meters per cubic centimeter. The volume surface area can be converted to the specific surface, in square meters per gram, by dividing by the 35 density of the material. Portland cement density is considered 3.15 grams per cubic centimeter as in

ANSI/ASTM standard C 204-78a; slag density is considered 2.92 grams per cubic centimeter.

The procedure for analyzing slag samples in the Microtrac^R is as follows: Two grams of slag are dispersed in reagent grade alcohol (denatured ethanol) until the equilibrium uncalibrated sample volume (dV) is reached without the presence of air bubbles. Each sample is analyzed in three 30 second runs, the results are then averaged.

The particle size data resulting from a triplicate set of Microtrac measurements are a table of volume percent of sample larger than a given size, diameter, in micrometers. The Microtrac^R instrument also furnishes the following summary data, employing these abbreviations:

	dV	Uncalibrated Sample Volume
-	%10	Microns at 10th percentile
	%50	Microns at 50th percentile
	%90	Microns at 90th percentile
20	MV	Volume Mean Diameter
•	CS	Calculated Surface Area (m^2/cc)
	STD. DEV.	Standard Deviation
	ма	Area Mean Diameter

Table I presents the particle size data for typical examples of the slag of this invention (M1) and the cement of this invention (M3). In Table II is a summary of the particle size data obtained on a number of samples.

Particle Size Distribution, Table I.

30	Channel	Percent	Larger Than
	Upper Limit	Sample Ml	Sample M3
	(um)		
	31	0.0	0.0
	22	0.0	1.1
35	16	0.0	3.9
	11	0.0	7.4
	7.8	1.5	14.9

	•		-	•		
			-	12 -		
	5.5		;	8.4	27.6	
	3.9		19	9.4	42.8	
	2.8		3	4.1	57.0	
	1.9		5	3.9	72.6	
5	1.4		7	6.0	87.8	
	0.9		9	2.9	97.8	
	0.9		10	0.0	100.0	
		Summar	y of Micro	otrac Data,	Table II	
		8	courser	% courser		
10	Sample	Code t	han llum	than 7.8um	<u>cs</u>	cm ² /gm
	Ml	4/17	0.0	1.8	3.901	13200
	Ml	5/8	0.0	0.8	4.035	13900
	Ml	5/8	0.0	0.9	3.710	12700
	Ml	8/15	0.0	1.5	3.176	10900
15	Ml	8/15	0.0	1.8	3.065	10500
	Ml	12/5	0.0	0.0	4.391	15100
	Ml	12/5	0.0	0.2	3.981	13600
	Ml	12/5	0.0	0.4	3.856	13200
	мз	5/8	7.4	14.9	2.263	7180
20	мз	5/8	7.5	14.7	2.321	7370
	M5	4/17	2.3	10.9	2.570	8480
	M5	5/8	1.3	7.5	2.876	9490
	FS	4/17	2.3	8.6	2.572	8810
	FS	8/15	2.0	5.5	2.886	9880
25	FS	8/15	1.4	4.4	2.762	9460
	os	8/15	27.7	39.8	1.683	5760
	OP I	12/29	50.3	59.5	1.219	3870
	OP III	12/29	40.0	49.2	1.667	5290
	Symbol	ls used	in tables	and example	25:	
30	Ml = ul	ltrafine	slag of	this		•
	in	nvention		01	P I=type	I portland
	M3 = u	ltrafine	cement	OP II	I=type II	[I portland
	M5 = u	ltrafine	commerci	al slag/cem	ent (80/2	20)
	FS = f	ine slag	•			
35	os = 0	ordinary	slag			

Set Time and Other Physical Measurements

Set time is determined by the following

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procedure, employing the formulations given in the tables.

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- 1. Water is added to a plastic cup.
- NaOH and Na₂CO₃ are added as required and dissolved.
- 3. Dispersant is added and dissolved.
- 4. Cementitious material is added and agitated for 10 seconds by a high speed mixer.
- 5. The cup is tilted periodically while the CM surface is observed. When there is no flow or gross motion of the CM surface while the cup is tilted to 45 degrees the time is noted; this is the set time.

other observations are also made on the sample. A qualitative estimate of the viscosity is made after the gross air entrained by the agitation has left but well before the sample has set.

Estimated viscosities are: medium - approximates midrange motor oil, high - approximates glycerine, low water viscosity; all at room temperature. The color
of the sample is observed; in some formulations this
is a quick indication of hardening. Particularly at
high water ratios, settling of the CM is observed; the
percent of the total volume present as clear surface
liquid is reported as percent settlement.

Hardness development is monitored by measurement of the compressive strength and is reported along with the time after sample preparation. The strength measurement employs a hand-held penetrometer (Soiltest CL-700 Pocket Penetrometer, Soiltest Inc., Chicago, Illinois) and follows the manufacturers instructions. The penetrometer is held by its handle with the piston at right angles to the CM surface. With steady pressure the piston is pushed into the CM to the depth of the calibration groove

about 1/4 form the end of the piston. The compressive strength is read on the scale of the instrument.

- Examples 1 and 2 show the effects of water ratio, NaOH content and Na₂CO₃ content on the set time and other properties. It is seen that formulations with NaOH have lower viscosities than those with Na₂CO₃, thus promote ease of permeation by the grout, but have higher set times.
- M1, M3 and M5 are as defined in Summary of Microtrac Data, Table II above. N2 is the dispersant, at 43% solids aqueous, sodium napthalene sulfonate formaldehyde polymer (CAS No. 9084-06-4).

When present sodium silicate

15 (SiO₂/Na₂O=3.22; employed as 37.8% aqueous) is further diluted with the second water charge to make the second component of a two component system. The two components are then mixed and the timing started.

 H_3PO_4 , orthophosphoric acid, is 85% acid 20 (aqueous).

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Example 1. Accelerators

			1:1 Wate	er/CM	Ratio					
	Formulation	<u>a</u> A	В	С	D	E	F	G	H	I
	Water (gms)	50	50	50	50	50	50	50	50	50
5	NaOH "	1	2	4				0.5	1	2
	Na ₂ CO ₃ "				1	2	4	0.5	1	2
	N2 (43%) "	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5
	Ml (gms)	50	50	50	50	50	50	50	50	50
	Results		•							
10	Viscosity	low	low	low	med	med	high	low	low	med
	minutes	15	13	12	10	9	9	7	6	5
	Set time (ır) 24	8	4.8	4.7	1.7	0.6	9	4.7	1.2
	$kg/cm^{2(1)}$	>4.5	>4.5	>4.5	0.25	>4.5	>4.5	>4.5	>4.5	3.0
	hours	33	21	21	21	21	21	21	21	4.7
15	$kg/cm^{2(1)}$				>4.5				;	>4.5
	hours				34					7.8
	Color	1b	1b	1b	1b	1b	dm	lb	din	dm
	hours	46	21	21	21	21	21	21	21	47
		(1) Unc	onfined	Compi	ressiv	e Str	ength			
20		Col	or code:	:						
	2	lb lig	ht blue							

mb medium blue

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Example 2. Accelerators

2:1 Water/CM Ratio

	Formulation	A	В	С	D	E	F	G	H	I
	Water (gms)	100	100	100	100	100	100	100	100	100
5	NaOH "	2	4	6				1	2	3
	Na ₂ CO ₃ "				2	4	6	1	2	3
	N2 (43%) "	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5
	Ml (gms)	50	50	50	50	50	50	50	50	50
	<u>Results</u>								•	
10	Viscosity	low	low	low	med	med	high	low	med	high
	hours	1.4	1.4	1.4	1.4	1.4	1.4	1.4	1.4	1.4
	Settlement(%)	50	50	30	20	20	20	30	20	10
	Set time (hr)	8	6	4	8	6	4.5	9	4.5	2.1
	$kg/cm^{2(1)}$	0.5	2.0	2.3	soft	soft	0.5	4.0	0.25	5 4.0
15	hours	18	14	14	20	20	18	14	75	7.5
	$kg/cm^{2(1)}$	2.3	3.7	3.7	>4.5	>4.5	>4.5	>4.5	>4.5	>4.5
	hours	20	20	20	33	33	20	20	10	10
	$kg/cm^{2}(1)$	>4.5	>4.5	>4.5						
	hours	72	72	72						
20	Color	mb	lb	lb	1b	1b	lb	lb	1b	lb
	hours	72	14	14	33	33	18	14	10	10

(1) Unconfined Compressive Strength

Color code:

mb medium blue

25 lb light blue

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	Ex	amr	ole	3.
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	Compar	rison o	f Set I	<u>'imes</u>	for M	11, M3	, and	<u> M5</u>	
	Formulation	A	В	С	D	E	F	. G	H
	Water (gms)	50	50	50	50	100	100	100	100
5	N2 (43%								
	solids) "	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5
	Ml (gms)	50			40	50			40
	мз "		50		10		50		10
	м5 "			50				50	
10	Results								
	Viscosity	low	high	low	low	low	med	med	low
	minutes	15	10	8	5	60	5 5	50	45
	Settlement((%) 15	10	5	5	10	15	20	15
	minutes		75		67	55	50	41	36
15	Set time (h	rs) -	0.8	1.3	0.8	3.3	2.7	1.7	8.3
	$kg/cm^{2(1)}$	soft	0.7	4.0	3.5	soft	4.0	0.3	0.3
	hours	23	9	23	23	46	23	22	22
	$kg/cm^{2(1)}$	1.3	3.0	>4.5	>4.5	>4.5	>4.5	>4.5	>4.5
	hours	48	12	48	46	129	46	45	46
20	$kg/cm^{2(1)}$	>4.5	>4.5						
	hours	94	23						
	Color	w	' dg	db	db	w	dg	db	þ
	hours	94	96	96	96	47	46	45	45
	1	(1) Unc	onfined	d Comp	ressi	ve St	rengt	h	
25		Col	or code	: :					
	7	w=white							
	C	ig=dark	gray						

db=dark blue

b=blue

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Example 4.

		Set	Time (of Sil	icate	/CM				
	<u>Formulation</u>	A	В	С	D	E	F	G	H	I
	Water (gms)	50	50	60	60	60	60	60	60	60
5	N2 (43%) "	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5
	Ml (gms)	50		18	16	20			10	15
	мз "			2	4		20		10	5
	M5 "		50					20		
	Water "			40	40	40	40	40	40	40
10	Silicate SiO ₂	$/Na_2O$								
	(3.22,	37.8%) (gms) 38	38	38	38	38	38	38
	Results									
	Set time(min)	22hr	8hr	20	6	6hr	0.5	1.7	2	3 .
	Settlement(%)	10	20	0	0	50	0	0	0	0
15	hours	14	14			34				
	$kg/cm^{2(1)}$	soft	soft	soft	soft	soft	0.75	0.7	1.0	0.2
		14	14	45	8	34	1	8	4	4
	$kg/cm^{2(1)}$ >	4.5	4.5	soft		soft	3	1.2	1.8	1.3
		82	82	94	70.	93	4	22	19	19
20	$kg/cm^2(1)$			soft	>4.5	soft	>4.5	>4.5	>4.5	>4.5
	hours		. }	183	172	183	20	70	82	82
	Color	¥	b	W	đ	W	g	g	đ	g
		82	82	94	70	93	20	22	34	34

(1) Unconfined Compressive Strength

25 Color code:

w=white

b=blue

g=gray

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Exa	mpl	e	5.

	Set Time of Silicate/CM										
	Formulation	A	В	С	, D	E	F	G	H	I	
	Water (gms)	50	50	50	50	50	50	60	60	60	
5	N2 (43%) "	0.5	0.5	0.3	0.3	0.5	0.5	0.2	0.2	0.2	
	NaOH (gms)	2	•	0.5		1.0		0.8			
	Na ₂ CO ₃ "		2		0.5		1.0				
	Ml "	50	50	25	25	50	50	20			
	M5 "								20		
10	мз "									20	
	water "	40	40	20	20	40	40	40	40	40	
	H ₃ PO ₄ (85%)(9	ms)						•	0.4	0.4	
	Silicate as	in									
	Ex. 4 (gms)	38	38	19	19	38	38	3.8	38	38	
15	<u>Results</u>										
	Viscosity	med	high	low	low	low	med	low	low	low	
	Set Time(hr)	0.15	0.25	0.7	1.5	0.33	0.5	4.7	0.16		
	$kg/cm^{2(1)}$	0.3	s	S	s	S	vs	vs	0.1	0.2	
	hours	0.66	15	15	15	15	14	18	4	0.5	
20	$kg/cm^{2(1)}$	1.5	s	S	s	1.2	vs		0.2	0.5	
	hours	15	40	24	24	24	24		18	4	
	$kg/cm^{2(1)}$	4.5	1	S	s	s	VS			0.7	
	hours	24		39	39	39	38			17	
	(1) Unce	nfined	Comp	ressi	ve St:	rengti	'n			

(1) Unconfined Compressive Strength Qualitative hardness code:

s=soft

vs=very soft

Example 6.

Permeation Test

The apparatus consists of:

- (1) a cell filled with fine sand, closed at each end by a stopper having a single central hole,
- (2) a pressurized grout supply tankconnected to the bottom of the cell by tubing and
 - (3) a delivering system consisting of a

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tube, connected to the top of the cell, leading to a graduated cylinder collector vessel. The cell is a 5 cm I.D., 15 cm long glass cylinder having a sand-retaining assembly at each end.

Each assembly consists of:

- (1) a 70 mesh per inch polyester sieve cloth for retaining the sand,
- (2) a close-fitting ca 5 cm 0.D. flanged disk with 12 holes 1/16 inch in diameter therethru,
- (3) a rubber stopper to seal the cylinder and to bear on the flange of the disk thereby creating a small plenum chamber between the disk and the stopper and
- (4) a metal end plate having a central hole to accommodate the tube passing therethru and edge-holes to accommodate threaded rods. Wing nuts on the threaded rods tighten down on the end plates thus forming a frame which clamps the assembly together so it can be pressurized without leaking.

Permeation Test Procedure

- 25 1. Press a rubber stopper into an end of the glass cylinder and then put a perforated disk and a sieve cloth into the tube.
 - 2. Weigh enough Evanston beach sand for one sample.
 - 3. Pour one third of the sand into the glass cylinder.
 - 4. Place a solid metal cylinder as a weight (about 320 grams) on the top of the sand, and apply a vibrator in the sand or against the glass.
- 5. Repeat steps 3 and 4 twice: then place a sieve cloth and a perforated disk on the top of the sand.
 - 6. Press a rubber stopper on the other end of the

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tube and assemble the frame.

- 7. Conn ct the supply and delivery tubes and apply pressure to water which is in the tank in order to saturate the sample.
- 5 8. Disconnect the tube connected to the bottom of the sample and let the water drain out by gravity.

 Record the volume of the water drained from the tube and label it as VI. This is the void volume in the sample before the injection of grout.
- 10 9. Prepare the grout and pour it into the emptied tank.
 - 10. Apply 2 psi air pressure to the tank and collect the effluent from the top of the sample. Time the accumulated volumes of effluent and label them as V2, V3 etc.
 - 11. The volume of slurry permeating the sample is V1 plus V2, V3, or V4 respectively.

	•			•					
		Perme	<u>eation</u>	Test_	Result:	<u> </u>			
	Formulation Programme 1	A	В	C	D	E	F	G	H
20	CM	Ml	M5	Ml	FS	os	Ml	Ml	Ml
	Water:CM	1:1.	2:1	2:1	2:1	2:1	2:1	2:1	2:1
	Dispersant	N2	N2	N2	N2	N2	N2	HSR	HSR
	% solids								
	on CM	0.43	0.43	0.43	0.43	0.43	0.43	0.6	0.02
25	Sand Wt (g)	337	355	332	336	326	336	356	344
	Sand Ht (cm)	9.7	10.1	9.0	9.8	9.7	9.9	10.4	10.1
	Sand d. (g/cc)	1.68	1.7	1.78	1.66	1.62	1.64	1.65	1.64
	V1 (cc)	13	13	14	11	13	12	14	13
	V2+V1 (cc)	55	51	99	62	30	112	92	60
30	minutes	1	1	1	1	1	1	1	1
	V3+V1 (cc)	81	101	204	93	38	249	162	65*
	minutes	5	5	5	5	5	10	10	5*
	V4+V1 (CC)	89*	123	326	122	62			
	minutes	10*	12	20	20	20			

The test results showed the differences in permeation among the grouts consisting of two parts

*flow stopped

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water to one part cementitious material, by weight, employing the N2 dispersant at 0.43% solids on cement. Data are also furnished (1) at a 1:1 ratio with water and (2) employing another dispersant, Lubricon HSR (American Admixtures, Inc., Chicago, IL 60646) a salt of Hydroxylated carboxylic acids.

Example 7.

Permeability of Grouted Formation

A confined column of Evanston beach sand is 10 grouted with each of the following grout formulations, in parts by weight:

		A	В.	С
	Ml	100	100	100
	N2, solids basis	0.4	0.4	0.4
15	NaOH	1	5	10 .
	Water	200	200	200

The permeability of the sand after the grout $\bar{}$ is hardened is very low being in the range 10^{-7} to 10^{-8} centimeters per second.

The Evanston beach sand is a fine sand having the following particle size distribution.

US Standard Sieve: 40 50 70 100 200 Wt. percent finer: 99 91 36 4 1

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CLAIMS

- 1. A composition comprising: (1) slag as a first cementitious material, the slag having a specific surface area greater than 9500 square centimeters per gram and less than three weight percent of particles larger than 7.8 micrometers and (2) an equal or greater weight of water.
- 2. The composition of claim 1, additionally comprising an effective amount of dispersant, to disperse the particles of slag, and the composition comprising up to 12 parts water per part slag, by weight.
- The composition of claim 2, additionally comprising up to 15% NaOH, Na_2CO_3 or a mixture thereof based on the dry weight of the slag, and the slag having a specific surface area greater than 11000 square centimeters per gram and less than $1\frac{1}{2}$ %, by weight, of particles larger than 7.8 micrometers.
- The composition of claim 3, consisting essentially of one part slag and, based on the weight of the slag, 1.5 to 7 parts water, up to about 2% dispersant, and up to about 10% NaOH, Na₂CO₃ or a mixture thereof.
- 5. The composition of claim 4, the slag having a specific surface area greater than 13000 square centimeters per gram with no particles over 7.8 micrometers in diameter and, based on the weight of slag, about 2 to 3 parts water, about 0.2% to 0.5% dispersant and about 3% to 8% NaOH, Na₂CO₃ or a mixture thereof.
- The composition of claim 1, additionally comprising a cement as a second cementitious material, the cement having a specific surface area greater than 7000 square centimeters per gram and less than 16 weight percent of particles larger than 7.8 micrometers, the slag being at least 40% of the total cementitious material by weight; the composition comprising up to 12 parts of water per part cementitious material by weight and an effective amount of a dispersant to disperse the cementitious material.
- 7. The composition of claim 6, the slag having a specific surface area greater than 11000 square centimeters per gram with less than 11% by weight of particles larger than

- 7.8 micrometers, the cement having a specific surface area greater than 8000 square centimeters per gram and less than 7% by weight of particles larger than 7.8 micrometers, slag being at least 60% of the antitious material by weight, and additionally comprising NaOH, Na₂CO₃ or a mixture of these being up to 15% of the slag plus up to 4% of the cement by weight.
- 8. The composition of claim 7, comprising, per part cementitious material c eight, from 1.5 to 7 parts of water and up to 2% dispersant.
- 9. The composition of claim 8, consisting essentially of: slag having a specific surface area greater than 13000 square centimeters per gram and no particles larger than 7.8 micrometers; cement having a specific surface area greater than 9000 square centimeters per gram and less than 3% by weight of particles larger than 7.8 micrometers; the slag being at least 75% by weight of the cementitious material; two to three parts of water per part of cementitious material by weight; 0.2% to 0.5% dispersant by weight based on the cementitious material; and Na₂CO₃, NaOH or a mixture of these being 3% to 8% of the slag plus zero to 2% of the cement by weight.
- 10. The composition of claim 2, additionally comprising up to 150% alkali silicate based on the weight of slag.
- 11. The composition of claim 10, the slag having a specific surface area greater than 11000 square centimeters per gram and less than 1½%, by weight, of particles larger than 7.8 micrometers, and, based on the weight of the slag, up to 2% dispersant and up to 10% orthophosphate ion.
- 12. The composition of claim 11, consisting essentially of slag having a specific surface area greater than 13000 square centimeters per gram and no particles larger than 7.8 micrometers; 1.5 to 7 parts of water per part of slag by weight; and, based on the weight of slag, 30 to 75% alkali silicate, 0.2% to 0.5% dispersant and 1 to 5% orthophosphate ion.
- 13. The composition of claim 10, additionally comprising

- a cement as a second cementitious material, the cement having a specific surface area greater than 7000 square centimeters per gram and less than 16 weight percent of particles larger than 7.8 micrometers, the slag being at least 40% of the cementitious material by weight.
- 14. The composition of claim 11, comprising: slag having a specific surface area greater than 11000 square centimeters per gram and less than 11% of particles larger than 7.8 micrometers, cement having a specific surface area greater than 8000 square centimeters per gram and less than 7% of particles larger than 7.8 micrometers, and based on the weight of cementitious material, up to 2% dispersant and up to 10% orthophosphate ion.
- The composition of claim 14, consisting essentially of: slag having a specific surface area greater than 13000 square centimeters per gram and no particles larger than 7.8 micrometers; cement having a specific surface area greater than 9000 square centimeters per gram and less than 3% by weight of particles larger than 7.8 micrometers; the slag being at least 75% by weight of the cementitious material; 1.5 to 7 parts of water per part of cementitious material by weight; and based on the weight of cementitious material, 30 to 75% alkali silicate, 0.2% to 0.5% dispersant and 1 to 5% orthophosphate ion.
- 16. The composition of claim 11, additionally comprising NaOH, Na_2CO_3 or a mixture of these, being up to 10% of the slag by weight.
- 17. The composition of claim 14, additionally comprising NaOH, Na_2CO_3 or a mixture of these, being up to 10% of the slag by weight.
- 18. A method of grouting comprising preparing the composition of claim 1 and injecting the composition into a formation.
- 19. A formation grouted by the composition of claim 1.

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INTERNATIONAL SEARCH REPORT

International Application No. PCT/US88/00197

I. CLASSIFICATION OF SUBJECT MATTER (if several classification symbols apply, indicate all) 6								
According to International Patent Classification (IPC) or to both National Classification and IPC								
IPC (4) CO4B 7/14								
U.S. CL. 106/117								
II. FIELDS	SEARCH	IED						
Minimum Documentation Searched 7								
Classification	n System		Classification Symbols					
U.S. C	L.							
•	U.S. CL. 52/744							
U.S. CI	U.S. CL 106/117							
Documentation Searched other than Minimum Documentation								
to the Extent that such Documents are Included in the Fields Searched 8								
III. DOCUMENTS CONSIDERED TO BE RELEVANT 9								
Category •		on of Document, 11 with indication, where app	ropriate, of the relevant passages 12	Relevant to Claim No. 13.				
Calcyony			2-5,10-19					
Y	US,A,	4402,749 (Hall et al), 6 Sept	emper 1903	2-3,10-13				
		olumn 5, lines 60-63; col. 2,	nnes 10-20,					
	colu m	n 3, line 5-10.						
		•						
X	IT C A	, 4443260, (Miyoshi, et al),	17 April 1984	1,6-9				
Y	00, 2	bstract; column 2, line 46; c	olumn 5. line 61.					
	Exam		· · · · · · · · · · · · · · · · · · ·	2-5,10-19				
	LAGII	· ·	•					
Y	115 4	, 4,415,364, (Naito, et al), 15	November 1983	2-5, 10-19				
-	See A	bstract.						
	300	. 0002 440						
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	00,, 7,11, 1,000, 1,0							
	L	of sited decomposite 10	"T" later document published after t	he international filing date				
 Special categories of cited documents: 10 "T" later document published after the international filling date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the 								
considered to be of particular relevance invention								
"E" earlier document but published on or after the international "X" document of particular relevance; the cannot be considered novel or cannot								
"I." document which may throw doubts on priority claim(s) or involve an inventive step								
citation or other special reason (as specified) cannot be considered to involve an inventive step when the								
"O" doc	"O" document referring to an oral disclosure, use, exhibition or other means "O" document is combined with one or more other such document is combined with one or more other such documents, such combination being obvious to a person skilled							
"P" document published prior to the international filing date but								
later than the priority date claimed								
IV. CERTIFICATION								
Date of the Actual Completion of the International Search Oate of Mailing of this International Search Report								
2 5 MAY 1988								
19 April 1988 International Searching Authority Signature of Authorized Officer								
International Searching Authority Signature of Authority Signature of Authority								
ISA/U	JS		Andrew Griffis					